

Summary of cosmid clones isolated from M. echinospora genomic library.	S_b resistance $(\mu g m L^{-1})^c$	0.5	6.6	0.5	0.5	0.5	0.5	0.1	< 0.01	0.05	0.04	< 0.01
	deoxy sugar gene	N.D. ^d	$N.D.^d$	+	+	+	N.D. ^d	+	+	+	+	1
	type I PKS genes b type II PKS genes b deoxy sugar genes b	N.D. ^d	$N.D.^d$	+	+	+	N.D. ^d	+	1	+	1	1
	type I PKS genes ^b	N.D. ^d	N.D. ^d	+	+	+	$N.D.^d$	+	1	+	ŀ	1
	clone ^a	3a	4a	4b	10a	13a	16a	56	58	09	99	puc18/pBluescript ^e

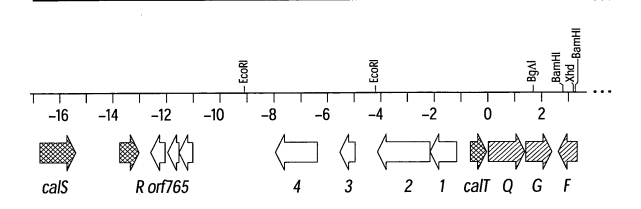
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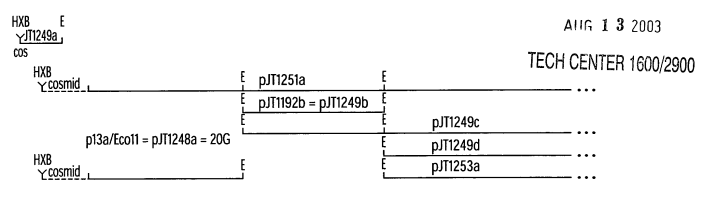
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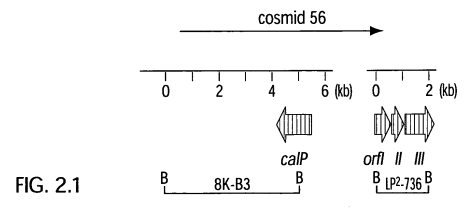




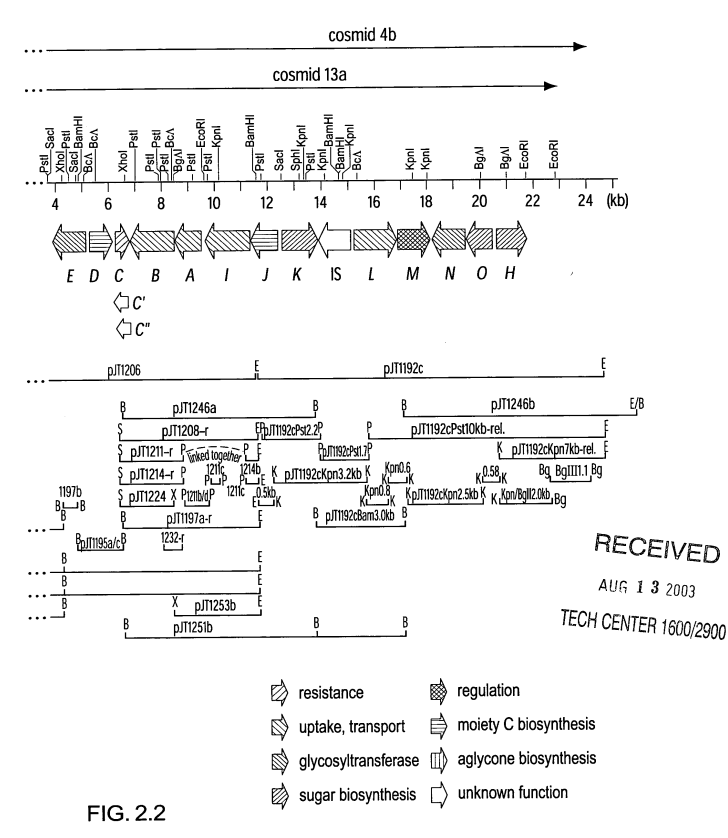
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putative polypeptide	number of amino acids	proposed function or sequence similarity detected	probability	start/stop codons	best match	
CalA	328	membrane transporter (ATP-binding)	5.4×10 ⁻¹²⁴	ATG/TGA	DrrA ⁹⁷	
CalB	561	membrane transporter	5.5×10 ⁻⁷⁰	ATG/TGA	DrrB ⁹⁷	
CalC	181	calicheamicin resistance protein	confirmed	ATG/TGA		
CalD	263	O-methyltransferase	1.1×10^{-99}	ATG/TGA	AveBVII ⁹⁸	
CalE	420	glycosyltransferase	4.7×10^{-30}	GTG/TAG	EryCII ⁹⁹	
CalF	245	N,N-dimethyltransferase	1.5×10^{-78}	ATG/TGA	DesVI ¹⁰⁰	
CalG	990	TDP-D-glucose 4,6-dehydratase	confirmed	GTG/TAG		
CalH	338	Perosamine synthetase	confirmed	GTG/TGA		
Call	568	Dipeptide transporter	1.7×10^{-24}	GTG/TGA	DciAE	
CalJ	332	O-methyltransferase	1.0×10^{-37}	ATG/TGA	DmpM	
CalK	440	L-cysteine/cystine C-S-lyase	1.6×10^{-28}	GTG/TGA	C-DES	
CalL	562	Oligopeptide transporter protein	9.5×10 ⁻¹⁴	ATG/TGA	OppA	
CalM	416	Regulatory protein	70	GTG/TGA		
CalN	398	Glycosyltransferase	3.4×10^{-79}	ATG/TGA	Ole1	
CalO	331	Hexopyranosyl-2,3-reductase	4.9×10^{-139}	ATG/TGA	EryBII	
CalP	(179)	Desaturase	5.7×10 ⁻⁷	/TGA	CrtI	
CalQ	453	UDP-D-glucose 6-dehydrogenase	confirmed	GTG/TGA	0.05.05.03	
CalR	282	Transcriptional regulator	6.7×10 ⁻¹¹	ATG/TGA	SC5C7.03	
CalS	1113	P ₄₅₀ oxidase	2.9×10 ⁻⁶⁶	GTG/TGA	BioI	
CalT	432	oxygenase/halogenase	2.0×10^{-62}	GTG/TAA	PCZA361.20	
CalU	377	glycosyltransferase	2.0×10^{-53}	ATG/TGA	SnogE/D	
CalV	125	β-keto-acyl synthase III	2.0×10^{-65}	ATG/TGA	SC4A9	
CalW	(449)	cytochrome P450	1.0x10 ⁻⁹¹	GTG/TGA	CYP105B1	
CalX		TDP-4-keto-6-deoxy-L-hexose 2,3-dehydratas	1.0×10^{-22}	/TGA	MtmV	
6MSAS	(198)	orsellenic acid synthase	6.5x10 ⁻⁷⁶	ATG/	AviM	
ActI	(207)	polyketide cyclase	3.0x10 ⁻⁶⁶	/TGA	Curr	~
ActII	136	polyketide cyclase	5.0x10 ⁻⁵³	ATG/TGA	SCIID A	76-
ActIII	(308)	polyketide synthase	8.6x10 ⁻¹⁴⁸	GTG/	Pms1	PECENEL 13 2003 1600200
<i>orf</i> l	322	unknown		ATG/TGA	* AU	
orf2	654	unknown	20 10-13	ATG/TGA	Soft of	10 0
orf3	373	integrase	3.0×10^{-13}	ATG/TGA GTG/TAA	Portella	2000
orf4	521	chromosome partitioning protein	$3.3x10^{-10}$	ATG/TGA	raiA 1//	b _ ~
orf5	175	unknown		ATG/TGA	•	16000
orf6	139	unknown		GTG/TGA		1 75m
orf7	187	unknown	3.0x10 ⁻⁶⁶	ATG/TGA	KorSA	
orf8	266	regulatory protein	1.5x10 ⁻⁷	ATG/TGA	SC4C6.24c	
Orf	127	hydroxylase	1.3X10	GTG/TGA	50-100.2-10	
OrfII	248	unknown	3.3x10 ⁻⁹⁰	GTG/TGA	SCA32	i
OrfIII	298	hydroxylase	5.3x10 ⁻⁴³	GTG/TGA	SC9C7.25	
<i>Orf</i> IV	363	unknown	2.9×10^{-37}	GTG/TGA	SCF55	
OrfV	288	aminotransferase		GTG/TGA	SCA32	
OrfVI	1012	glu-ammonia-ligase adenylytransferase	8.0x10 ⁻⁶³	GTG/TGA	SCF43A.25c	
OrfVII	236	Methyltransferase	8.9x10 ⁻⁹	GTG/TGA	SCA32	
OrfVIII	441 479	Integral membrane protein	1.1x10 ⁻²¹	ATG/TGA	MLB268	
OrfIX	478 504	Integral membrane protein	5.5x10 ⁻²⁰	GTG/TGA	B1496.F1.14	
OrfX	504	Membrane protein Immunity resistance protein	1.1x10 ⁻⁹	ATG/TGA	TFXG	
OrfXI	251	insertional element	5.7×10 ⁻¹⁶⁸	1110/1011	IS <i>1136</i> ¹¹¹	
IS-elemen	t 1209 bp	inscritonal element	3.1710			J

FIG. 3



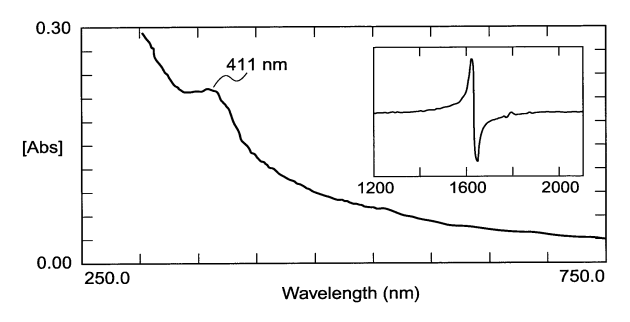


FIG. 4A

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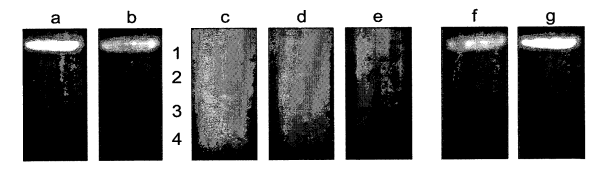
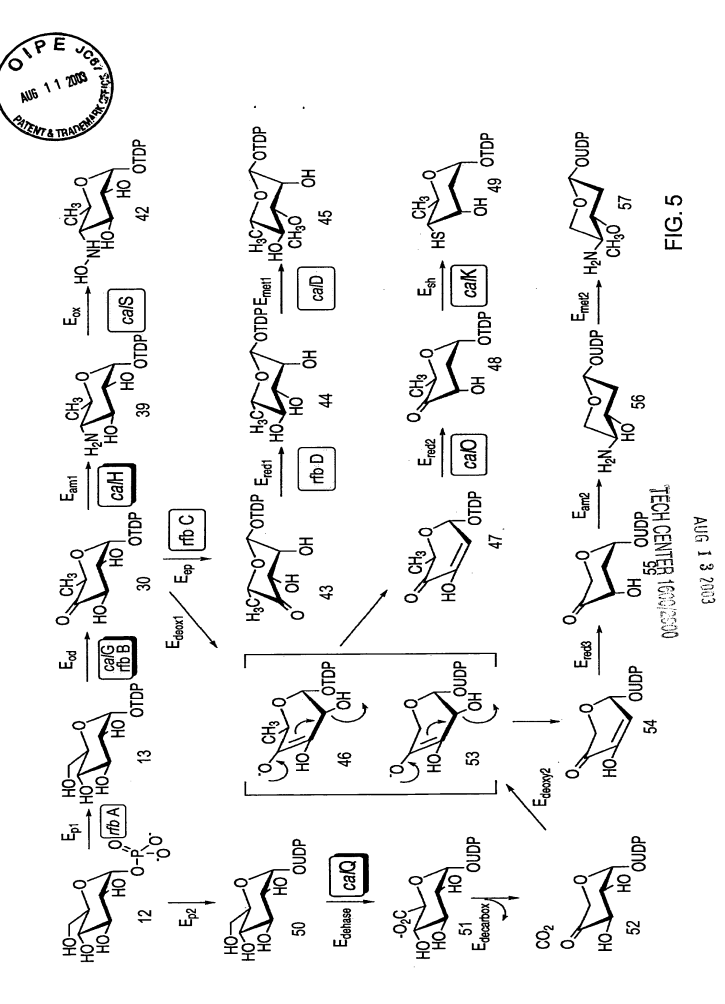
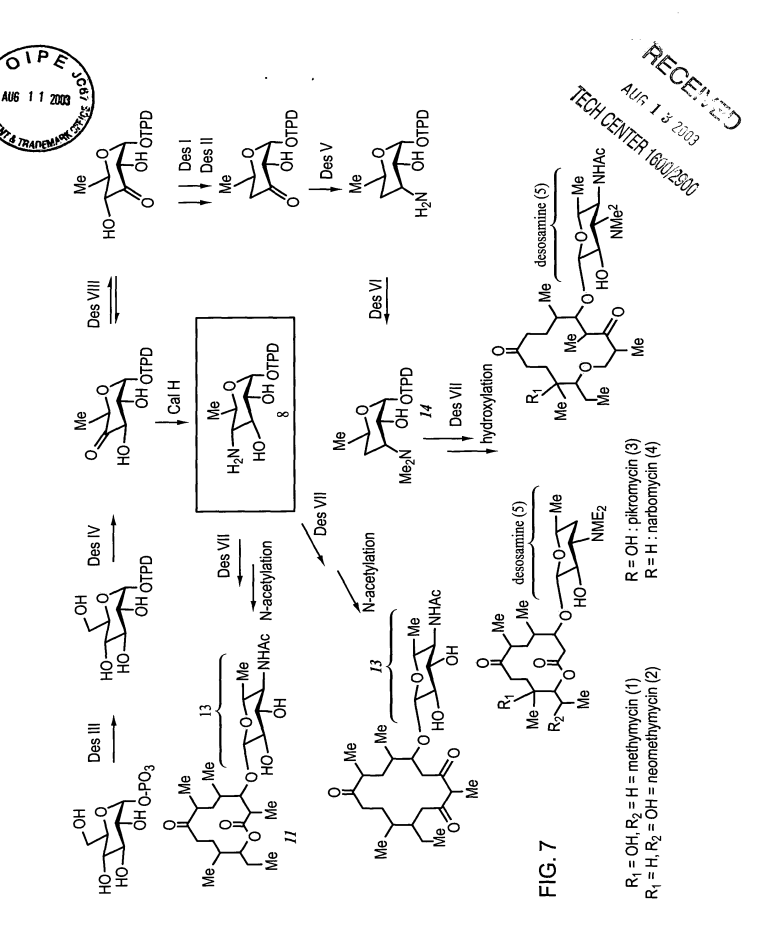
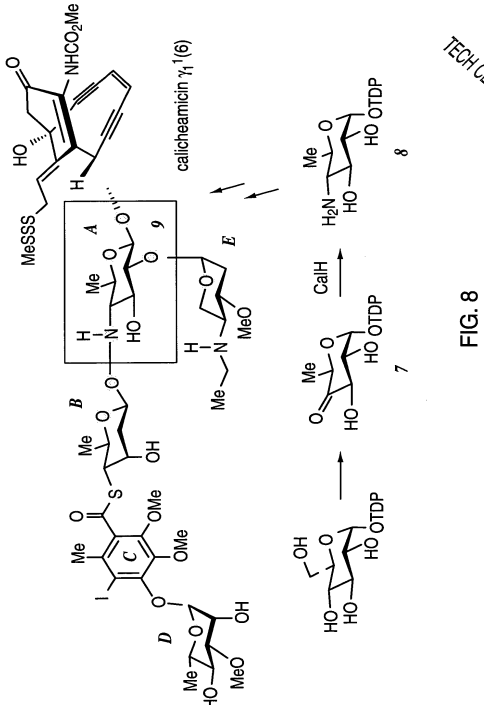


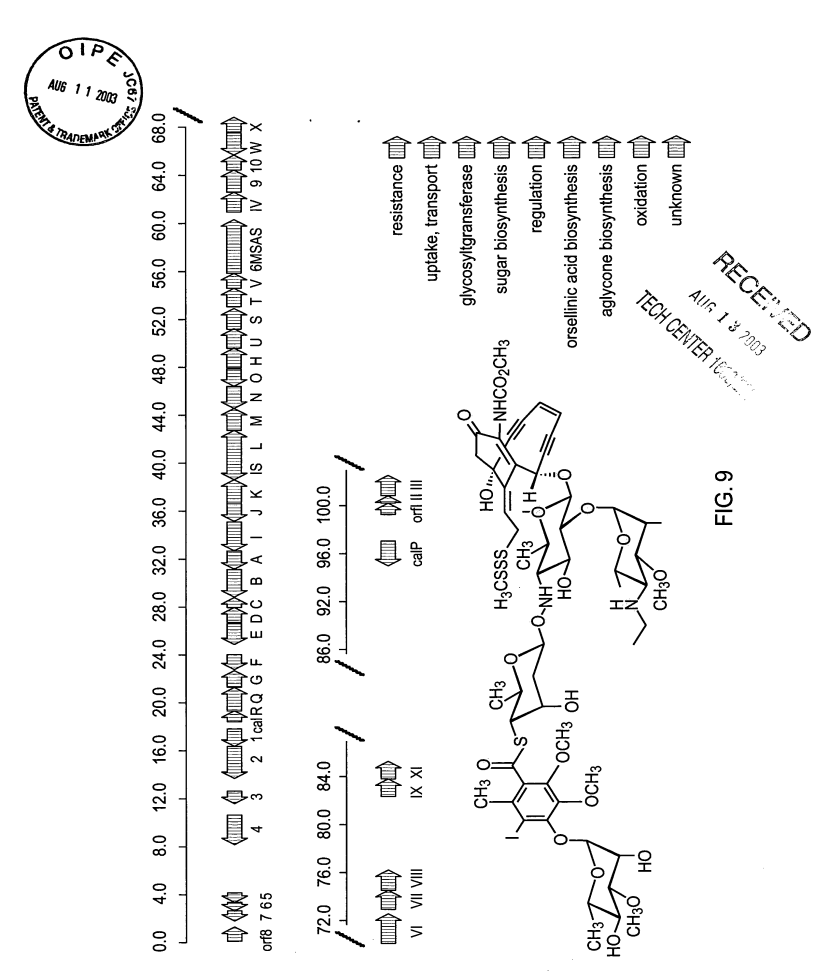
FIG. 4B

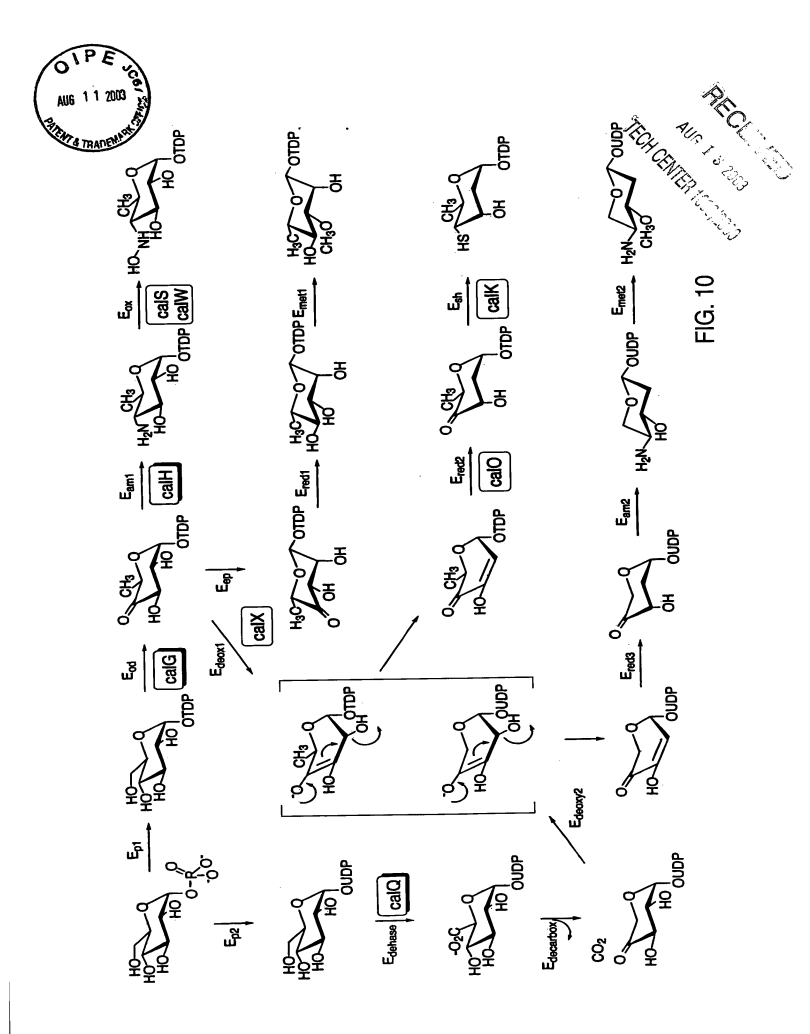








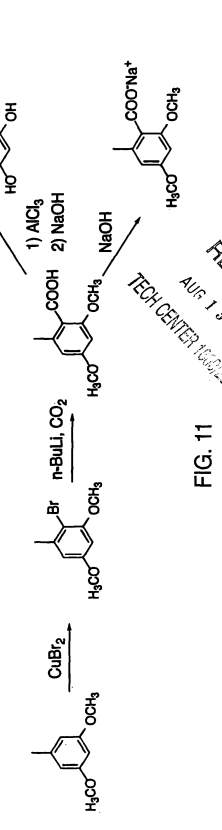






The Aryltetrasaccharide Unit (a type I PKS product):

Synthesis of Putative Substrates:



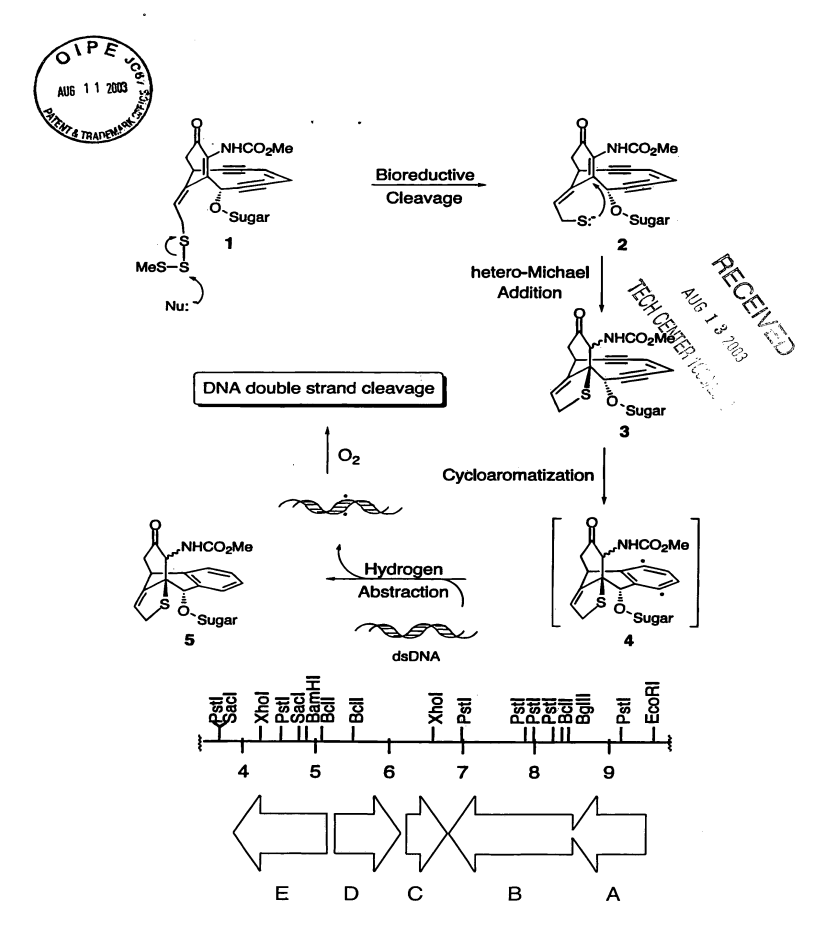
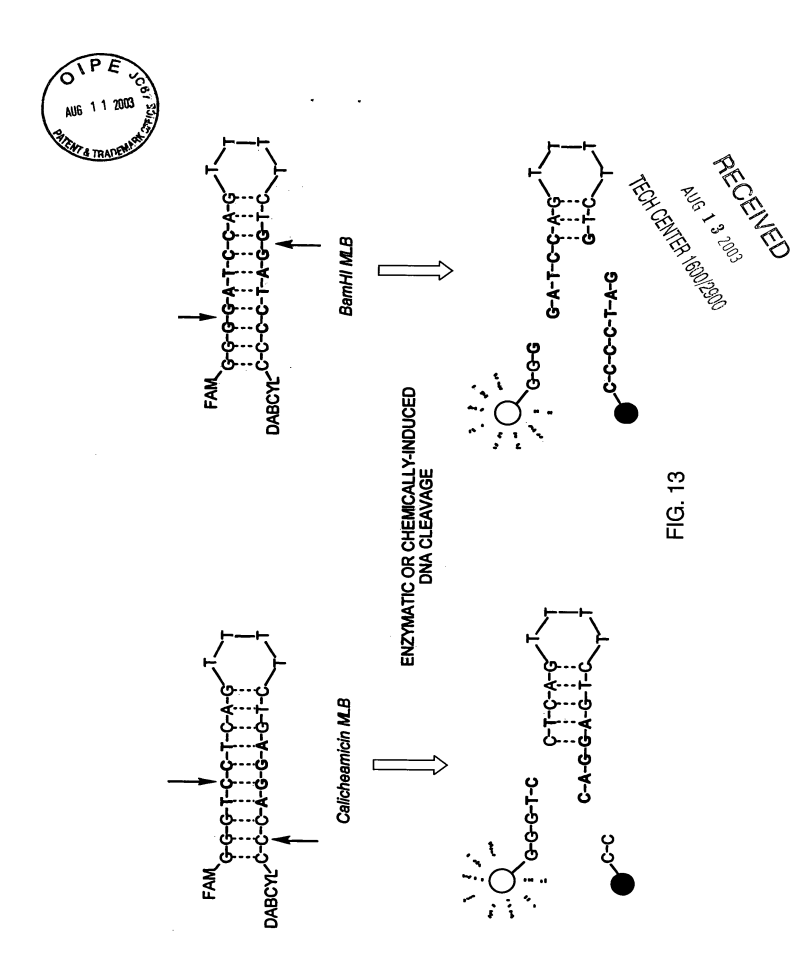
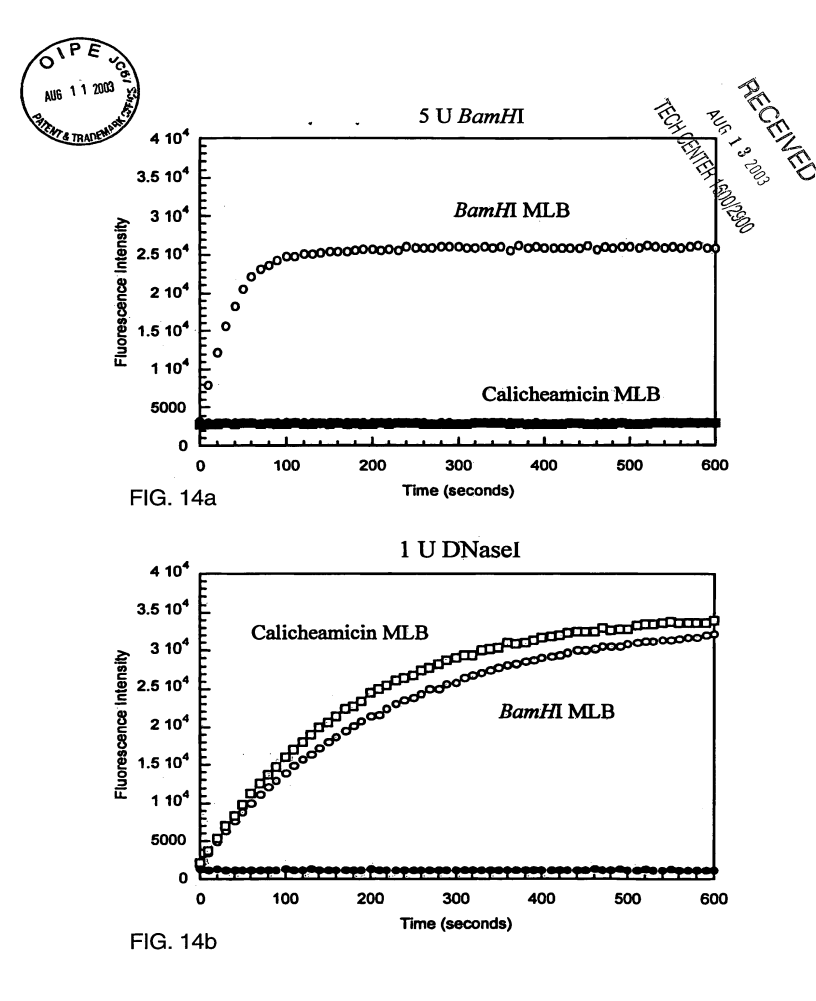


FIG. 12







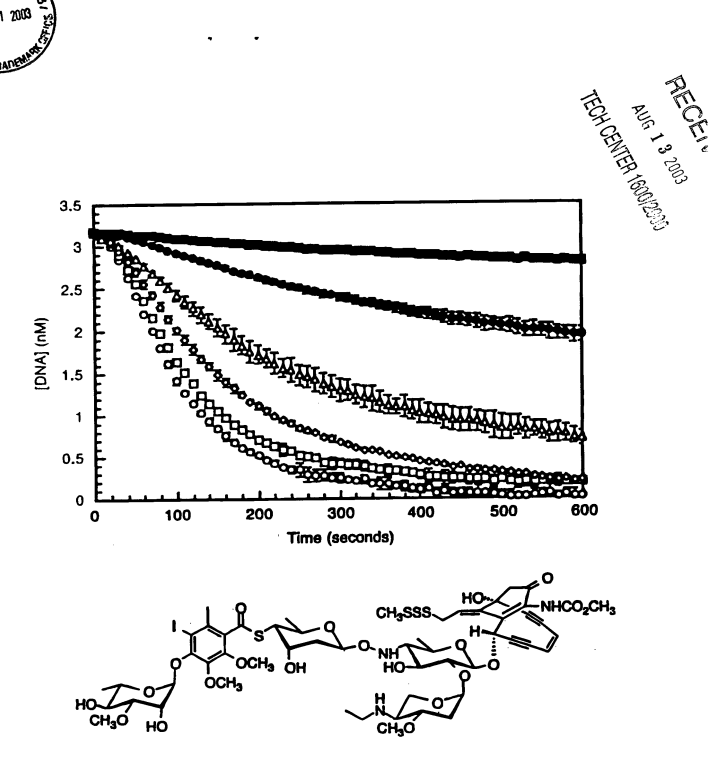


FIG. 15a



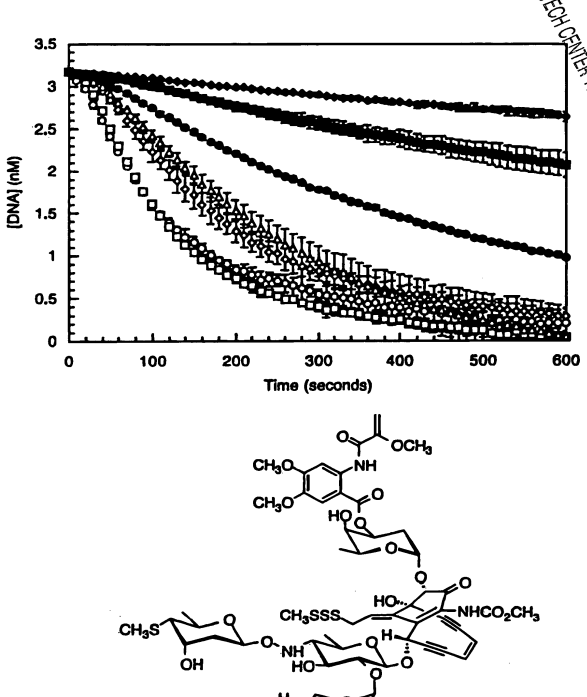


FIG. 15b





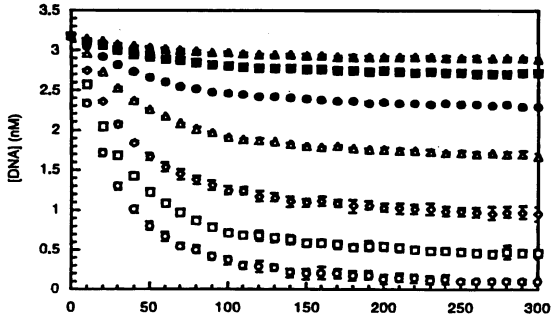


FIG. 16a



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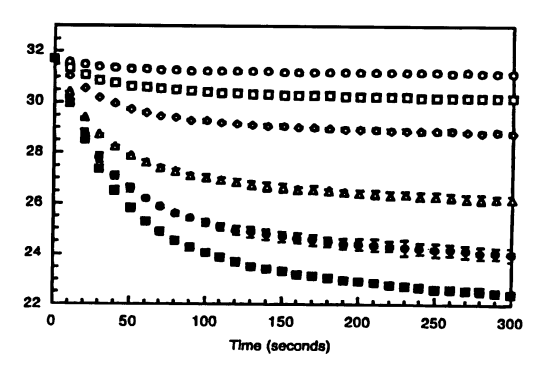
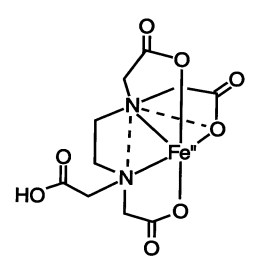


FIG.16c





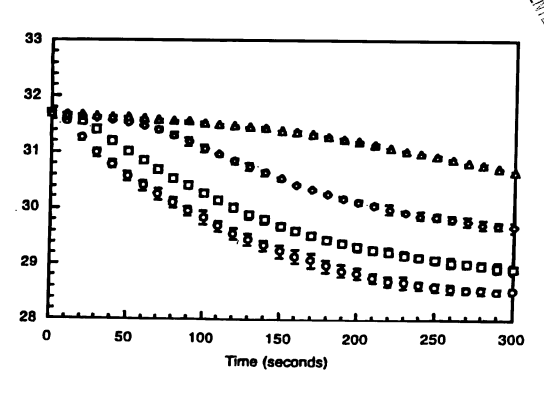


FIG. 16d

